PATENT SPECIFICATION

NO DRAWINGS.

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COMPLETE SPECIFICATION.

Therapeutic Preparations Containing 7-Substituted Theophylline Derivatives.

We, LES LABORATOIRES DAUSSE, a French Body Corporate, of 4 rue Aubriot, Paris, France, do hereby declare the invention, for which we pray that a patent may be granted 5 to us, and the method by which it is to be performed, to be particularly described in and by the following statement:-

This invention relates to therapeutic preparations containing 7-substituted theo-

phylline derivatives.

According to the present invention there is provided a therapeutic composition of matter comprising (a) a purine component having a musculotropic action which is a water-soluble, 15 7-substituted theophylline derivative, such as $7 - \beta$ - hydroxy - ethyl theophylline, $7 - \beta - \gamma$ - dihydroxyropyl theophylline and salts of theophylline - 7 - ethanoic acid; and (b) an adrenergic component which is the hydrochloride of 1 - (3:4 - dihydroxyphenyl)-2-methylamino-1-propanol.

It has been found that a medicinal synergy exists between the hydrochloride of 1-(3:4dihydroxyphenyl) - 2 - methylamino - 1 - propanol and the purine components as herein-

before defined.

The potentiated bronchodilatory effect-obtained by the administration of the composition - containing 1 - (3:4 - dihydroxy, phenyl) - 2 - methylamino - 1 - propanol, acting by means of an adrenergic mechanism, and the above-defined purine components, of which the action is mainly musculotropic, are particularly useful in the treatment of bronchial dyspnea and more especially asthma.

This potentiation has been shown by the method of recording the tonus of the bronchi of the guinea pig as described by Halpern

(Arch. Int. Pharmacodyn. et Therap., 1942,

68, 339).

The minimum active doses A and P of the adrenergic component and of the purine component on acetylcholinic bronchospasm having been determined, doses A1 and P1 of each of these components, lower than the doses A and P respectively, are chosen, and it is found that they have no action on the bronchospasm produced by the injection of acetylcholine.

Continuing the experiment, there are simultaneously administered to the guinea pig the dose A^1 of adrenergic component and the dose P1 of purine component, and it is found that this association is capable of 55 inhibiting and sometimes even suppressing the bronchospasm produced by acetylcholine, the latter being employed in the same dose throughout the experiment.

Thus, the simultaneous administration of an ineffective does A1 of the hydrochloride of 1 - (3:4 - dihydroxyphenyl) - 2 - methylamino-1-propanol and of an ineffective dose P1 of a purine component, or of a mixture of purine components, produces by mutual potentiation an unexpected bronchodilatory effect, since it is greater than the sum of the effects peculiar to each of the constituents of the composition.

The new synergic compositions have many advantages.

In the first place they permit of obtaining a considerable bronchodilatory effect by utilising only small quantities of the substances constituting the composition. Thus, the desired therapeutic effect can be fully obtained despite the reduction of the posology of each of the constituents, which results in a

50

	lowering of the toxicity without a diminution	(2) 1 - (3:4 - Dihydroxyphe-
	of the activity.	nyl) - 2 - methylamino - 1-
	For example, it is known that adrenergic substances, of which 1-(3:4-dihydroxphenyl)-	propanol hydrochloride 0.025 g.
5	2 - methylamino - 1 - propanol hydrochloride	7 - β - γ - Dihydroxypropyl
	is one, produce fairly frequently tachycardia	theophylline 4 g. 70
	and signs of central excitation which result in	Reducing solvent q.s 50 ml.
	trembling of the extremities, notably of the hands, and insomnia.	To but the state of the state o
0	The synergic action of the purine bases	In both cases, the reducing solvent em- ployed is a solution of the following com-
•	makes it possible to reduce the dose of 1-	position:—
	(3:4 - dihydroxyphenyl) - 2 - methylamino-	Lamasa.
	1-propanol and to reduce to a very consider-	Sodium bisulphite solution 2.5 ml. 75
	able extent, or to eliminate, the secondary	Disadism substitution of the
5	effects in question.	
	Since the purine bases also have central stimulating effects characterised essentially	Distilled water q.s 1000 ml.
	by insomnia, it is desirable to add to the	It is to be noted that these solutions can be
	synergic compositions of the present inven-	distributed in 1 ml. or 2 ml. ampoules, so
)	tion a quantity of a drug which is a barbituric	that there are obtained either ampoules con- 80
	derivative. Butobarbital or butylethyl-	taining ½ mg. or ampoules containing 1 mg. of
	malonylurea has proved particularly desirable from this standpoint.	1 - (3:4 - dihydroxyphenyl) - 2 - methyl-
	The compositions may comprise in addi-	amino-1-propanol hydrochloride. These ampoules (preferably those of 1 ml.
,	tion one or more other purine substances	containing only ½ mg. of 1-(3: 4-dihydroxy- 85
	selected from the ophylline, the ophylline ethy.	phenyl) - 2 - methylamino - 1 - propanol hy-
	lenediamine and caffeine. The new compositions are of value in the	drochloride) may be used for shallow sub-
	treatment of respiratory troubles of bronchial	cutaneous or intramuscular injections.
	or pulmonary origin, of asthma, of pulmonary	
	emphysema, of chronic bronchitis, of pul-	Example II.
	monary sclerosis, of chronic catarrh of the	Aqueous solution for atomisation:— 90
	respiratory passages and of silicosis.	(1) Ampoule A
	The purine component and the adrenergic component may be associated with an	
	excipient for suppositories, an aqueous	1 - (3 : 4 - Dihydroxyphe- nyl) - 2 - methylamino - 1-
	excipient for parenteral administration, an	propanol hydrochloride . 0.01 g.
	aqueous excipient for administration by the	Monosodium sulphite solu- 95
	aerial route or an excipient for oral admini- stration.	tion 0.003 ml.
	When the composition is used in an aqueous	Distilled water q.s 1 ml.
	medium, it is desirable to take account of the	
	tendency of the diphenol, which is 1-(3: 4-di-	Ammoule D
	hydroxyphenyl) - 2 - methylamino - 1 - pro-	Ampoule B
	panol, to oxidise in the presence of com- pounds having an alkaline reaction. It is	7 - β - γ - Dihydroxypropyl theophylline 0.375 g. 100
:	therefore important to avoid the choice of a	Distilled meeting 10 1
٠	theophylline derivative having an alkaline	Distinct water q.s 10 ml.
	reaction and it is preferred that there should	(T)
	be included in the aqueous medium an anti-	The contents of the two ampoules are
,	oxidant or a reducing agent which is acceptable from the pharmacological riomagist	mixed and the mixture administered in aerosol form by discharge from a pressurised
•	able from the pharmacological viewpoint, for example sodium bisulphite or sodium	container. 105
1	formaldehyde sulphoxylate.	(2) The following single solution composi-
	Examples of pharmaceutical forms of the	tions may also be adopted, the reducing sol-
	compositions of the present immedian and the	vent being that which is specified for solutions
	compositions of the present invention are the	
	following :—	intended for parenteral administration.
1	following : EXAMPLE I.	intended for parenteral administration.
1	ollowing :— EXAMPLE I. Parenteral Administration :—	intended for parenteral administration. 1 - (3:4 - dihydroxyphenyl) - 2- 110
1	following : EXAMPLE I.	1 - (3:4 - dihydroxyphenyl) - 2- methylamino - 1 - propanol
1	EXAMPLE I. Parenteral Administration:— (1) 1 - (3:4 - Dihydroxyphenyl) - 2 - methylamino 1- propanol hydrochloride 0.025 g.	1 - (3:4 - dihydroxyphenyl) - 2- 110 methylamino - 1 - propanol hydrochloride 0.01 g.
1	EXAMPLE I. Parenteral Administration:— (1) 1 - (3:4 - Dihydroxyphenyl) - 2 - methylamino 1- propanol hydrochloride 0.025 g. 7 - β - γ - Dihydroxypropyl	 1 - (3:4 - dihydroxyphenyl) - 2- 110 methylamino - 1 - propanol hydrochloride 0.01 g. 7 - β - γ - Dihydroxypropyl theo-
1	EXAMPLE I. Parenteral Administration:— (1) 1 - (3:4 - Dihydroxyphenyl) - 2 - methylamino 1- propanol hydrochloride 0.025 g.	1 - (3:4 - dihydroxyphenyl) - 2- 110 methylamino - 1 - propanol hydrochloride 0.01 g.

		•		
	Example III.		Lac varnish 0.005 g.	
	Suppositories:—		Absorbent powder . 0.005 g.	
	(Î) For adults :—		Taleum 0.02 g.	55
	1 - (3:4 - Dihydroxyphe-		Crystallised sugar . 0.13 g.	
5	nyl) - 2 - methylamino - 1-		Erythrosin traces	
	propanol hydrochloride	0.005 g.	Carnauba wax traces	
	7 - β - γ - Ďihydroxypropyl			
	theophylline	0.30 g.	WHAT WE CLAIM IS:—	
	Sodium hydrosulphite	0.002 g.	and the Continue	60
10	Eutectic mixture of glycer-		1. A therapeutic composition of matter	60
	ides of fatty acids of natural		comprising (a) a purine component having a	
	vegetable origin (m.p. +		musculotropic action which is a water-	
	35° C.)	1.655 g.	soluble 7-substituted theophylline deriva-	
	·	-	tive; and (b) an adrenergic component	۵,
	(2) For infants:—		which is the hydrochloride of 1-(3:4-di-	65
15	1 - (3:4 - Dihydroxyphe-		hydroxyphenyl) - 2 - methylamino - 1 - pro-	
	nyl) - 2 - methylamino - 1		panol.	
	propanol hydrochloride	$0.0015\mathrm{g}$.	2. A composition according to Claim 1	
	$\tilde{7}$ - $\tilde{\beta}$ - γ - $\tilde{\mathbf{D}}$ ihydroxypropyl		wherein the theophylline derivative is 7-β-	70
	theophylline	0.085 g.	hydroxyethyl theophylline, 7-β-γ-dihydroxy-	70
20	Sodium hydrosulphite	$0.0019 \mathrm{g}.$	propyl theophylline or a salt of theophylline-	
	Cochineal carmine	0.0004 g.	7-ethanoic acid.	
	Eutectic mixture of glycer-		3. A composition according to Claim 1 or	
	ides of fatty acids of natural		2 wherein the purine component and the	75
	vegetable origin (m.p. +		adrenergic component are associated with an	75
25	35° C.)	1.800 g.	excipient for suppositories, an aqueous	
			excipient for parenteral administration, an	
	(3) With butobarbital:—		aqueous excipient for administration by the	
	1 - (3:4 - Dihydroxyphe-		aerial route or an excipient for oral admini-	80
	nyl) - 2 - methylamino - 1-	0.005	stration.	00
	propanol hydrochloride	0.005 g.	4. A composition according to Claim 3	
30	$7 - \beta - \gamma - Dihydroxypropyl$	0.00	wherein the excipient contains a pharma-	
	theophylline	0.30 g.	cologically acceptable antioxidant or reducing	•
	Butobarbital	0.05 g.	agent. 5. A composition according to any of	85
	Sodium hydrosulphite	0.002 g.	Claims 1—4 which contains in addition a	
	Eutectic mixture of glycer-		drug which is a barbituric acid derivative.	
35	ides of fatty acids of natural		6. A composition according to Claim 5	
	vegetable origin (m.p. +	1 COE ~	which contains butobarbital.	
	35° C.)	1.605 g.	7. A composition according to any of	90
	Thursday, TV		Claims 1—6 which further contains one or	
	Example IV.		more other purine substances selected from	
••	Tablets:—		theophylline, theophylline ethylenediamine	
40	7 - β - γ - Dihydroxy-		and caffeine.	
	propyl theophyl-		8. A therapeutic composition of matter	95
	line 0.04 g.)		according to Claim 1 substantially as herein-	
	Caffeine 0.06 g.		before described with reference to any of the	
	1 - (3 : 4 - Dihydroxy-		foregoing specific examples.	
45	phenyl - 2 - methyl-		1010Gong abouts arm-base	
	amino - 1 - propa-	Nucleus:	J. A. KEMP & CO.,	
	nol hydrochloride 0.01 g.	0.20 g.	Chartered Patent Agents,	
	Icing sugar 0.02 g. Maize starch 0.01 g.	0.20 g.		
20			14 South Square,	
50	2 222		Gray's Inn,	
			London, W.C.1.	
	Talcum 0.0455 g. J		-	
				

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